Document AN3 Appl. No. 09/956,980



WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau

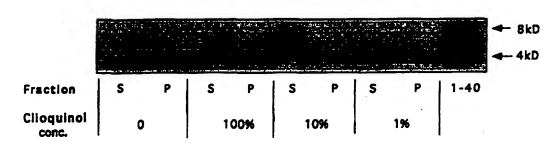


INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶ : A61K 31/47	A1	(11) International Publication Number: WO 98/06403
		(43) International Publication Date: 19 February 1998 (19.02.98)
(21) International Application Number: PCT/IB (22) International Filing Date: 8 August 1997 (BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE,
(30) Priority Data:		LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ,
960100286 13 August 1996 (13.08.96) (71) Applicant (for all designated States except US,		BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN,
GEROLYMATOS S.A. [GR/GR]; 13 Asklipiou, 65 Kryoneri Attikis (GR).	GR-1	ML, MR, NE, SN, TD, TG).
(72) Inventor; and (75) Inventor/Applicant (for US only): GEROLYMATO: otis, Nikolas [GR/GR]; 13 Asklipiou, GR-145 65 Attikis (GR).	S, Pana Kryon	Published With international search report. With amended claims.
(74) Agents: BAGGER-SØRENSEN, Birgitte et al.; Inte- Patent-Bureau, Høje Taastrup Boulevard 23, DK-26 trup (DK).	mation 530 Tai	ait is-

(54) Title: USE OF THE CHELATING AGENT CLIOQUINOL FOR THE MANUFACTURE OF A PHARMACEUTICAL COMPOSITION FOR THE TREATMENT OF ALZHEIMER'S DISEASE

Solubilisation of βAmyloid with Clioquinol



(57) Abstract

The use of the chelating agent clioquinol for the manufacture of a pharmaceutical composition for the treatment of Alzheimer's disease is disclosed.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM .	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	Prance	LU	Luxembourg	SN	Senegal
ΑŪ	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
١Z	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Paso	GR	Greece		Republic of Macedonia	TR	Turkey
3G	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
IJ	Benin	IB	Ireland	MN	Mongolia	UA	Ukraine
3R	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
3Y	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada _	ľΤ	lta ly	MX	Mexico	UZ.	Uzbekistan
F	Central African Republic	JP	Japan	NE	Niger	VN	Vict Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
H	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
1	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
M	Cameroon		Republic of Korea	PL	Poland		
N:	China	KR	Republic of Korea	PT	Portugal		
יטי	Cuba	KZ	Kazakstan	RO	Romania		
:Z	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
E	Germany	u	Liechtenstein	SD	Sudan		
K	Denmark	LK	Sri Lanka	SE	Sweden		
LE .	Estonia	LR	Liberia	SG	Singapore		

PCT/IB97/00983 WO 98/06403

USE OF THE CHELATING AGENT CLIOQUINOL FOR THE MANUFACTURE OF A PHARMA-CEUTICAL COMPOSITION FOR THE TREATMENT OF ALZHEIMER'S DISEASE

The present invention relates to the use of a chelating agent for the manufacture of a pharmaceutical composition for the treatment of Alzheimer's disease.

Alzheimer's disease is a chronic neurodegenerative disease prevalent primarily among elderly people.

The clinical diagnosis of Alzheimer's disease is supported by the existence and accumulation of amyloid deposits in the brain. The amyloid is primarily found in the terminal zones of neurons as morphologically heterogeneous deposits, also known as senile plaque.

15 The formation of senile plaque is related to the appearance of the symptoms and signs of the disease, including amnesia. After the formation of senile plaque, neurofibrillary tangles are produced in the neuronal bodies. The formation of neurofibrillary

20 tangles is related to the worsening of amnesia and of the other symptoms of dementia.

A major component of the amyloid deposits is a polypeptide referred to herein as Aβ (Amyloid-beta). Aβ is normally a soluble component of the cerebrospinal fluid where it is found in concentrations of about 3-5 nm. Aβ may have 39 to 43 amino acids, typically 40 amino acids, in the matured form and is derived as a proteolytic cleavage product from a cell surface protein called the amyloid precursor protein (APP)

30 (Kang et al. 1987). The normal function of $A\beta$ is not known at present but might be to form cation-selective channels across cell membranes (Kawahara et al., 1997).

The precipitation of synthetic Aβ has been shown to be caused by several environmental factors including low pH, high salt concentrations and the presence of metals, e.g. zinc, copper, and mercury (Bush et al.,

1995). It has been reported that $A\beta$ itself specifically and saturable binds zinc of a high affinity binding (K_D = 107 nM) at a molar ratio of 1:1 (zinc: $A\beta$) (Bush et al., 1994a). This binding takes place at physiological concentrations of zinc (Bush et al., 1994b).

There is a strong supposition that the removal of amyloid deposits from patients suffering from Alzheimer's disease will alleviate the symptoms of Alzheimer's disease. Therefore, several attempts have been 10 made to prepare a drug for the removal of amyloid deposits.

WO 93/10459 discloses a method for the treatment of Alzheimer's disease by administrating a zinc binding agent. As preferred compounds phytic acid, desferri15 oxamine, sodium citrate, EDTA, 1,2-diethyl-3-hydroxypyridin-4-one, and 1-hydroxyethyl-3-hydroxy-2-methylpyridin-4-one are mentioned.

DE 39 32 338 discloses the use of a chelator, such as 8-hydroxy-quinoline, for the treatment of Alzhei-20 mer's disease.

US 5373021 discloses disulfiram and its salts and analogues in as far as these can penetrate the blood-brain-barrier. The disclosed compounds may be used to reduce neurological damages caused by i.a. Alzheimer's disease.

The hitherto known compounds suggested for the treatment of Alzheimer's disease have several drawbacks, which has prevented their widespread use. E.g., most of the compounds are unable to penetrate the blood-brain-barrier and thus can hardly reach the areas in which the amyloid is deposited. Disulfiram, which may penetrate the blood-brain-barrier, has the drawback that it also is an alcohol deterrent.

The most extensively studied chelator is EDTA. 35 However, the chelating effect of EDTA is relatively

3

weak towards zinc and copper. Furthermore, EDTA cannot penetrate the blood-brain-barrier and is considered relatively toxic.

The object of the present invention is to provide

5 a new use of a known pharmaceutical compound for the
manufacture of a pharmaceutical composition for the
treatment of Alzheimer's disease, said new compound
having the ability to penetrate the blood-brain barrier, to effectively chelate heavy metals to prevent

10 the aggregation of amyloid, and to redissolve amyloid
deposits. This object is achieved by using clioquinol
for such a new use.

The term "treatment of Alzheimer's disease" used herein refers to the prevention, control and therapy of 15 Alzheimer's disease at any stage of its development.

Clioquinol has the chemical name 5-chloro-7-iodo-8-hydroxyquinoline and belongs to the group of hydroxyquinolines. Clioquinol has a known use as topical anti-infective agent. Especially, clioquinol has been used 20 for the treatment of amoebiasis and infectious diarrhoea. Clioquinol is only scarcely soluble in water within a pH range of 7-11. Outside this range suitable concentrations may be achieved.

The chelating ability of clioquinol is known for 25 Fe, Co, Ni and Zn (Kidani et al., 1974 and Tateishi et al., 1973). Using mass spectrophotometry the coordination number for clioquinol in case of Co(II), Ni(II), Cu(II) and Zn(II) is 2, whereas the coordination number for Fe(III) is 3. Reportedly, injected preparations of clioquinol have crossed the blood-brain-barrier, leaving concentrations thereof in the brain in the order of 20µl/ml when administrated at dosages of 10-20 mg/kg (Tateishi et al. 1975 and Tamura 1975). The concentration of clioquinol was also found to be high in

4

such areas of the brain like the hippocampus which is immediately affected by Alzheimer's disease.

Using microautoradiograpic techniques, clioquinol has been shown in monkeys to form zinc chelates in the 5 hippocampus. The Zn(II) chelates were mainly found in the terminal axodendritic boutons of the mossy fibres. Unconjugated clioquinol has an extremely rapid penetration when injected intravenously into the nervous system with practically no blood-brain-barrier (Shinaki, 1979).

Even though clioquinol had a known chelating and blood-brain-barrier penetrating ability it could not have been predicted that clioquinol also had the ability to redissolve zinc precipitated $A\beta$. According to the present invention the new use of clioquinol for the manufacture of a pharmaceutical composition for the treatment of Alzheimer's disease is based on this unexpected finding.

At present, it is believed that clioquinol and $A\beta$ 20 competitively chelate zinc and other heavy metals. Clioquinol is regarded as the strongest chelator and will, therefore, predominately capture the heavy metal ions. Thus, $A\beta$ from precipitated zinc- $A\beta$ will be resolubilized into the surrounding fluid because clioquinol will capture the zinc ions. The complex of clioquinol and zinc will penetrate the blood-brain-barrier and be cleared from the organism.

Since clioquinol is a relatively strong chelator, it may also chelate metal ions from enzymes or pros30 thetic groups. Therefore, it might be desirable to supplement trace metal ions or prosthetic groups to clioquinol treated patients, especially when establishing prolonged clioquinol treatments.

Vitamin ${\bf B}_{12}$ contains cobalt. Previous case studies 35 of subacute myelo-optic neuropathy (SMON) in Japanese

5

patients who were administrated clioquinol at higher dosages than recommended and for long periods indicate a connection between prolonged clioquinol treatment and vitamin B_{12} deficiency. In the present application the interaction between clioquinol and B_{12} is proved through in vitro and in vivo studies. Therefore, it may be beneficial to administrate vitamin B_{12} together with or subsequent to the administration of clioquinol in order to prevent B_{12} deficiency.

Clioquinol may be administrated in any appropriate amount in any suitable galenic formulation and following any regime of administration.

Preferably, the amount of daily administration will be from 10 mg to 750 mg clioquinol depending on the condition of the patient. A typically daily dosage is 100 mg. Alternatively, from 10 mg to 250 mg, preferable 100 mg clioquinol, three times daily, may be administrated. A daily dosage of up to 750 mg for a period of two weeks is considered without any risk of 20 neurotoxicity or other side effects.

For the prevention of the onset of the symptoms and signs of Alzheimer's disease, or for the delay of the symptoms and signs in the evolution of the disease, daily clioquinol dosages of 10 mg to 100 mg can be administrated for long periods, viz. up to ten years.

The actual administrated amount is to be decided by the supervising physician and may depend on multiple factors, such as, the age, condition, file history, etc. of the patient in question. The results presented 30 herein suggest that a "window" for the administrated amount of clioquinol exists, within which the best dissolving results are obtained. Such a window may be determined by the physician upon conducting routine experiments.

6

If clioquinol is going to be administrated for a prolonged period, clioquinol is preferably administrated intermittently. In a first period clioquinol may be administrated e.g. for one to three weeks, followed by a wash-out period, which may provide for restoration of any unwanted side effects of clioquinol. The duration of the wash-out period may be one to four weeks. During the wash-out period it is preferred to administrate B₁₂ and other prosthetic groups and/or trace metals. After the wash-out period the first period may be repeated. The long-term intermittent therapy will provide not only for the resolubilisation of zinc-Aβ aggregates but also for the prophylactic inhibition of the formation of zinc-Aβ aggregates.

The intermittent administration of clioquinol will also reduce the toxicity potential of the drug, which means that the treatment may be extended throughout the evolution of the disease.

The pharmaceutical composition comprising clio20 quinol may be any suitable galenical formulation for
enteral and parenteral administration aiming at delivering sufficient concentrations of clioquinol into the
brain. It is preferred to administrate clioquinol as
intravenous injections if it is desired to obtain high
25 brain concentrations rapidly. As it is more convenient
to administrate a pharmaceutical composition orally,
this way of administration may be used if it is not of
importance to rapidly obtain high brain concentrations.
The pharmaceutical composition may also be admini30 strated intradermally.

The pharmaceutical composition comprising clioquinol may comprise other active ingredients as well. Especially, the composition may contain trace metals or prosthetic groups such as vitamin B₁₂, and/or any other therapeutic agent that can be used concurrently for the

7

treatment of Alzheimer's disease to improve or alleviate the symptoms and signs of Alzheimer's disease, or to delay the onset of any of the symptoms and signs of Alzheimer's disease. Alternatively, the other active ingredient(s) may be administrated as separate pharmaceutical composition(s) together with the pharmaceutical composition comprising clioquinol.

According to the invention, the new use of clioquinol comprises:

- 1) the treatment of patients already diagnosed as having Alzheimer's disease at any clinical stage,
 - 2) the prevention of the disease in the patients with early or prodromal symptoms and signs, and/or
- 3) the delay of the onset or evolution or aggrava-15 tion or deterioration of the symptoms or signs of Alzheimer's disease.

In the following the present invention will be illustrated by means of examples, however, these examples are not to be considered as limitating for the 20 scope of the protection.

EXAMPLES

EXAMPLE 1

25

5.3 g of clioquinol was suspended with agitation in 200 ml of n-decane. The undissolved material was allowed to settle. Weighing the dried undissolved clioquinol after blowing off the decane indicated that only 2% of the clioquinol dissolved in the decane. 100 ml of the (light yellow) supernatant was agitated together with 100 ml of PBS pH 7.4 and the phases allowed to separate. The PBS (lower phase) was collected and filtered to remove the residue which formed at the phase interface upon extraction with the organic

8

solvent. Assuming that 2% of the clioquinol dissolved in the n-decane, and assuming that the partitioning coefficient is 1/1750 with PBS at 1:1 mixtures of decane/clioquinol, the concentration of clioquinol in 5 the PBS is 800 nM.

Brain specimens were obtained for which a histophatological diagnosis of Alzheimer's disese was confirmed. Duplicate 0.5 g specimens of frontal lobe neocortex were homogenised in 3 ml of the clioquinol/
10 PBS solution at 100%, 10% and 1% of the final PBS/clioquinol extract and PBS alone.

The homogenates were centrifuged at 150,000 xg for thirty minutes and the supernatants collected and held on ice (fraction "S"). The pellets were subjected to an identical homogenisation and centrifugation regime and the resulting supernatants again collected (fraction "P").

1 ml of each supernatant was treated with 200 μ l of ice cold 10% TCA to precipitate total protein in-20 cluding A β . The resulting pellet was washed once with 100% ethanol and resuspended in 100 μ l of TBS (tris 20 mM, NaCl 150 mM pH 7.4).

7.5 μ l of sample (S or P) was boiled for 5 min with an equal volume of tris-tricine sample buffer 25 containing 4% SDS and loaded onto a Novex pre-cast 10-20% tris-tricine gel followed by Western transfer onto nitrocellulose. Signal for A β was detected using mAb WO2 (raised against residues 5-16 of A β) and visualized using ECL. The sensitivity of the detection system is 30 5-10 pg.

To validate the TCA precipitation for A β 1 μ g of synthetic A β 1-40 was added to 1 ml of PBS containing 10% BSA and the solution was treated as above. Signal for A β was detected in the precipitated pellet but not in the supernatant.

9

The result is indicated in fig. 1.

As it might be deduced from fig. 1, the clioquinol was effective in promoting the solubilisation of $A\beta$ in the concentrations tested. Furthermore, the optimal concentration was found to be "10%", indicating that one of the aggregation forms of $A\beta$, presumable the dimer, is more soluble in PBS than others.

While data are shown only for one specimen in fig. 1, data for 19 other specimens all indicate the same 10 tendency, viz. that clioquinol is effective in promoting the solubisation of $A\beta$.

Example 2

In this example the chelating ability of clioquinol and EDTA is compared.

Samples of 10 ng synthetic $A\beta$ were placed in microtitre wells and caused to aggregate by the addition of 25 μ M ZnCl. The aggregates were then transferred to a 0.2 μ m nylon membrane by filtration. The aggregates were washed with 200 μ l TBS alone, TBS containing 2 μ M EDTA, and TBS containing 2 μ M clioquinol. The membrane was fixed, probed with the anti- $A\beta$ monoclonal antibody 6E10 and developed for exposure to ECL-film. The transmittance of the ECL-film was measured and the relative signal strength calculated based on 100% for TBS alone. The relative signal strength was 66% for EDTA and 49% for clioquinol.

The results indicate that clioquinol is a better 30 chelator for zinc precipitated $A\beta$ than EDTA.

EXAMPLE 3

In this example the resolubilisating effect of 35 clioquinol is demonstrated.

10

A 2.5 μ M solution of A β in TBS at a pH of 7.4 was prepared. 95% of the A β was maintained in a soluble state. Addition of 30 μ M zinc resulted in precipitation of the soluble A β and only 43% was maintained in solution. The subsequent addition of 120 μ M clioquinol to the zinc precipitated A β resulted in an increase of soluble A β to 70%.

The results indicate that clioquinol is able to redissolve zinc precipitated $A\beta$.

10

EXAMPLE 4

In this example the influence of clioquinol on vitamin \mathbf{B}_{12} is studied in vitro using NMR spectroscopy.

Since clioquinol is practically insoluble in water within the pH range of 7-11, the study was conducted at pH 13. Three tubes were prepared. The first contained 1 mg clioquinol in 0.5 ml, the second contained 1.4 mg cyanocobalamine (B₁₂₎ in 0.5 ml, and the third contained 0.5 mg clioquinol + 0.7 mg cyanocobalamine in 0.5 ml (mol ratio 3:1).

The ¹H NMR spectra were recorded in a DRX 400 MHz spectrophotometer at 20°C. Comparison of the third spectra with the first and the second shows that some 25 of the resonances of cyanocobalamine are shifted and the same is observed for two of the resonances of clioquinol. The results suggest an interaction between clioquinol and cyanocobalamine.

30 EXAMPLE 5

In this example the influence of clioquinol on vitamin \mathbf{B}_{12} is studied in vivo.

Six weeks old male mice were pre-treated with 35 clioquinol for three days (50mg/kg/day). The mice were

11

divided into two groups, a control group and a group which was injected with [57Co]-cyanocobalamine. 48 hours after the injection the animals were sacrificed, and the brain, liver and kidney were dissected and counted in a gamma-counter as thousand cpm/g tissue (wet weight) ± SEM. The radioactivities in each of the groups are stated in Table 1 below:

Table 1

10	Treatment	Brain	Liver	Kidney
	Control	9.4 ± 0.9	97 ± 8	895 ± 207
	[⁵⁷ Co]-B ₁₂	8.4 ± 1.5	85 ± 21	252 ± 61

A comparison of the results show that there were 15 no significant changes in the amount of radioactive accumulation in the brain and the liver. A reduction in the amount of vitamin B_{12} trapped in the kidneys was apparent. It may be concluded that clioquinol has an influence on the concentration of B_{12} in some of the 20 mice organs.

REFERENCES

- Kang et al. (1987) Nature 325: 733-736.
- Kirschner et al. (1987) Proc. Natl. Acad. Sci. 267:
 6953-6957.
- Kawahara M.; Arispe N.; Kuroda Y.; Rojas E. (1997)
 Biophysical Journal 73/1, 67-75.
- Bush, A.I., Moir, R.D., Rosenkranz, K.M., and Tanzi, R.E. (1995) Science 268: 1921-1923.
- 10 Bush, A.I., Pettingell Jr., W.H., Paradis, M.D., and Tanzi, R.E. (1994a) J. Biol. Chem. 269: 12152-12158.
 - Bush, A.I., Pettingell Jr., W.H., Multhaup, G., Paradis, M.D., Vonsattel, J.P., Gusella, J.F., Beyre-
- uther, K., Masters, C.L., and Tanzi, R.E. (1994b) Science **265**: 1464-1467.
 - Esler, W.P., Stimson, E.R., Jennings, J.M., Ghilardi, J.R., Mantyh, P.W., and Maggio, J.E. (1996) Neurochem. 66: 723-732.
- - Shirilogi H., et al., (1979) Handbook of Clinical Neurology, North Holland Publishing Company, 141-198.
- 25 Tamura Z, (1975), Jap. J. med. Sci.Biol. Suppl 28: 69-77.

PATENT CLAIMS

- 1. Use of clioquinol for the manufacture of a pharmaceutical composition for the treatment of Alzheimer's disease.
- 2. Use according to claim 1, wherein the clioquinol is administrated in an amount of 10 to 250 mg one to three times daily.
- Use according to claim 1 or 2, wherein trace metals and/or prosthetic groups are administrated
 together with or subsequent to the administration of clioquinol.
 - 4. Use according to claim 3, wherein the prosthetic group is vitamin B_{12} .
- Use according to any of the proceeding claims,
 wherein the pharmaceutical composition comprising clioquinol is administrated intermittently.
- 6. Use according to claim 5, wherein the pharmaceutical composition comprising clioquinol is administrated for one to three weeks followed by a 20 wash-out period of one to four weeks.
 - 7. Use according to claim 6, wherein trace metals and/or prosthetic groups are administrated during the wash out period.
- 8. Use according to claim 7, wherein the prosthet- 25 ic group is vitamin B_{12} .
 - Use according to any of the preceding claims, wherein treatment is extended up to ten years.
- 10. Use according to any of the preceding claims, wherein the pharmaceutical composition comprising 30 clioquinol is formulated for oral administration.
 - 11. Use according to any of the claims 1 to 9, wherein the pharmaceutical composition is formulated for parenteral administration.

14

12. Use according to any of the claims 1 to 9, wherein the pharmaceutical composition is formulated for intradermal administration.

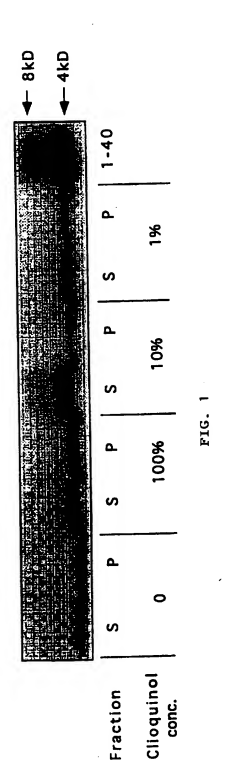
15

AMENDED CLAIMS

[received by the International Bureau on 15 January 1998 (15.01.98); new claims 13 and 14 added; remaining claims unchanged (1 page)]

- 12. Use according to any of the claims 1 to 9, wherein the pharmaceutical composition is formulated for intradermal administration.
- $\underline{5}$ 13. Pharmaceutical composition comprising clioquinol and vitamin B_{12} .
 - 14. Pharmaceutical composition according to claim 13, wherein the amount of clioquinol is 10 to 250 mg.

Solubilisation of β Amyloid with Clioquinol



International Application No PCT/IB 97/00983

A. CLASS	ification of subject matter A61K31/47		
According t	o International Patent Classification(IPC) or to both national clas	eification and IPC	
	SEARCHED		
Minimum di IPC 6	ocumentation searched (classification system followed by classif A61K	ication symbols)	
Documenta	tion searched other than minimum documentation to the extent th	nat such documents are included in the	fields searched
Electronic o	data base consulted during the international search (name of dat	a base and, where practical, search le	ms used)
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
A	WO 93 10459 A (UNIV MELBOURNE) cited in the application see abstract see page 6, line 13 - page 11, claims 15-21		1-12
A	DE 39 32 338 A (NMI NATURWISSENSCHAFTL U MEDIZ) 11 April 1991 cited in the application see the whole document		1-12
A	H. TJÄLVE: "THE ETIOLOGY OF S INVOLVE AN INTERACTION BETWEEN AND ENVIRONMENTAL METALS" MEDICAL HYPOTHESES, vol. 15, 1984, pages 293-299, XP002047206 see the whole document	MON MAY CLIOQUINOL	1-12
		-/	·
X Fur	ther documents are listed in the continuation of box C.	X Patent family members a	are listed in annex.
"A" docum	ategories of cited documents: ent defining the general state of the art which is not	cited to understand the principal	or the international filing date inflict with the application but siple or theory underlying the
"E" earlier filing o "L" docum which citatio "O" docum	dered to be of particular relevance document but published on or after the international data ent which may throw doubts on priority claim(s) or is cited to establish the publicationdate of another an or other special reason (as specified) ment referring to an oral disclosure, use, exhibition or means	involve an inventive step with "Y" document of particular relevant cannot be considered to invide the combined with ments, such combination be	or cannot be considered to en the document is taken alone
"P" docum later t	ent published prior to the international filling date but than the priority date claimed	in the art. "8" document member of the sar	
	actual completion of theinternational search	Date of mailing of the internal	шопа: зеагсп герогт ·
	7 November 1997	11/12/1997 Authorized officer	
Name and	mating address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswift Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Hoff, P	

International Application No PCT/IB 97/00983

A. CLASS IPC 6	A61K31/47		
According	to International Patent Classification (IPC) or to both national classifi	cation and IPC	<u> </u>
	S SEARCHED		
Minimum d	focumentation searched (classification system followed by classification $A61K$	tion symbols)	
Documente	ation searched other than minimum documentation to the extent that	such documents are included in the fields se	arched
Electronic	data base consulted during the international search (name of data b	ase and, where practical, search terms used)
C. DOCUM	IENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the re	elevant passages	Relevant to claim No.
A	WO 93 10459 A (UNIV MELBOURNE) 2 cited in the application see abstract see page 6, line 13 - page 11, 1 claims 15-21		1-12
A	DE 39 32 338 A (NMI NATURWISSENS MEDIZ) 11 April 1991 cited in the application see the whole document	CHAFTL U	1-12
A	H. TJÄLVE: "THE ETIOLOGY OF SMO INVOLVE AN INTERACTION BETWEEN O AND ENVIRONMENTAL METÄLS" MEDICAL HYPOTHESES, vol. 15, 1984, pages 293-299, XP002047206 see the whole document		1-12
X Furt	her documents are listed in the continuation of box C.	Patent family members are listed	in annex.
"A" docum: consid "E" earlier filing c "L" docume which citatio "O" docume other "P" docume later ti Date of the	ent defining the general state of the art which is not dered to be of particular relevance documents but published on or after the international date and which may throw doubts on priority claim(s) or is cited to establish the publicationdate of another in or other special reason (as specified) sent referring to an oral disclosure, use, exhibition or means ent published prior to the international filling date but han the priority date claimed actual completion of theinternational search. 7. November 1997	"T" later document published after the inte- or priority date and not in conflict with cled to understand the principle or th invention "X" document of particular relevance; the o- cannot be considered novel or canno- involve an inventive step when the do- "Y" document of particular relevance; the o- cannot be considered to involve an in- document is combined with one or m- ments, such combination being obvio- in the art. "8" document member of the same patent Date of mailing of the international asse-	the application but early underlying the claimed invention to complete to courner its taken alone claimed invention ventive step when the ore other such docu-us to a person skilled family
Name and r	making address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijaw(k Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3018	Authorized officer Hoff, P	

International Application No PCT/IB 97/00983

C /Ca=tl=	ntion) DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/IB 97/00983	
ategory *		Relevant to claim No.	
•	A.I. BUSH ET AL.: "MODULATION OF ABETA ADHESIVENESS AND SECRETASE SITE CLEAVAGE BY ZINC" THE JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 269, 1994, pages 12152-12158, XP002047207 cited in the application see the whole document	1-12	
A	J.E.F. REYNOLDS: "MARTINDALE, THE EXTRA PHARMACOPOEIA" 1993 , THE PHARMACEUTICAL PRESS , LONDON XP002047208 " Clioquinol " see page 511	1-12	
A	WO 95 31199 A (GEROLYMATOS P N SA ;LEGAKIS NICOLAOS (GR)) 23 November 1995 see abstract; claims	1-12	
			
	·		
		,	

3

information on patent family members

International Application No PCT/IB 97/00983

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 9310459 A	27-05-93	AU 669493 B CA 2123211 A	13-06-96 27-05-93
		EP 0613560 A JP 7503316 T	07-09-94 06-04-95
DE 3932338 A	11-04-91	NONE	
DE 3335330 V			
WO 9531199 A	23-11-95	AU 6580494 A EP 0713392 A	05-12-95 29-05-96